FILE 'HOME' ENTERED AT 13:37:59 ON 05 JUN 2006

=> file reg

Uploading C:\Program Files\Stnexp\Queries\10785070.str

```
chain nodes :
21 22 23 24 25 27
ring nodes :
1 2 3 4 5 6 7 8
                     9 10 11 12 13 14 15 16 17 18 19 20 28 29 30
31 32 33 34 35 36
                    37
chain bonds :
5-21 6-14 15-24 21-22 21-23 23-27 24-25
ring bonds :
1-2 1-6 2-3 2-7 3-4 3-10 4-5 5-6 7-8 8-9 9-10 11-12 11-16
                                                             11-17
12-20 13-14 14-15 15-16 17-18 18-19 19-20 28-29 28-32 29-30
                                                              30-31
33-34 33-37 34-35 35-36 36-37
exact/norm bonds :
21-22 21-23 23-27 24-25 24-41 28-29 28-32 29-30 30-31 31-32 33-34 33-37
34-35 35-36 36-37
exact bonds :
5-21 6-14 15-24
normalized bonds :
1-2 1-6 2-3 2-7 3-4 3-10 4-5 5-6 7-8 8-9 9-10 11-12 11-16 11-17 12-13
12-20 13-14 14-15 15-16 17-18 18-19 19-20
isolated ring systems :
containing 1 : 11 :
```

G1:H,CH3,Et,i-Pr,n-Bu,i-Bu,t-Bu

G2:OH, CN, [*1], [*2]

Match level :

Page 1

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 13:Atom 14:Atom 15:Atom 16:Atom 17:Atom 18:Atom 19:Atom 20:Atom 21:CLASS 22:CLASS 23:CLASS 24:CLASS 25:CLASS 27:CLASS 28:Atom 29:Atom 30:Atom 31:Atom 32:Atom 33:Atom 35:Atom 36:Atom 37:Atom 41:CLASS

L1STRUCTURE UPLOADED

=> d l1

L1 HAS NO ANSWERS

G1 H, Me, Et, i-Pr, n-Bu, i-Bu, t-Bu G2 OH, CN, [@1], [@2]

Structure attributes must be viewed using STN Express query preparation.

=> s l1 full 10 SEA SSS FUL L1

=> file ca

=> s 13

11 L3

=> d ibib abs hitstr 1-11

L4 ANSWER 1 OF 11 CA COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 144:191943 CA 144:191943 CA 144:191943 CA 145:191943 CA 145:191943 CA 146:191943 CA 146:19 CORPORATE SOURCE: Laboratoire de Stereochimie (CNRS UMR 7509), Universite Louis Pasteur (ECPM), Strasbourg, 67087, Fr. European Journal of Organic Chemistry (2005), (23), SOURCE: DUGY-5054 CODEN: EJOCFK; ISSN: 1434-193X Wiley-VCH Verlag GmbH & Co. KGAA Journal PUBLISHER: DOCUMENT TYPE: DERMY TYPE: Journal
JUNGE: English
1-Fluoro-2-(triethylsily))naphthalene and other 1-fluoronaphthalene
derive. bearing a metalation-resistant substituent at the 2-position
proved to be totally inert toward base attack. 3-Bromo-1fluoronaphthalene, readily prepared from a 2-bromo isomer by
deprotonation-triggered heavy helogen migration, was converted into
3,3'-dibromo-1,1'-difluoro-2,2'-binaphthyl [1] by consecutive treatment
with lithium diisopropylamide, copper(II) bromide and nitrobenzene. The
dilithiated intermediate generated from the atropisomer I by treatment
with 2 equivalent of butyllithium reacted with a variety of
trophiles to
afford products such as, diacid or bis(phosphane) derive. in high
yields. The latter compound was also obtained in a straightforward
ner from (4-fluoro-2-naphthyl)diphenylphosphine oxide. Unexpectedly, neither the 3,3'-dibromobinaphthyl I nor its 3,3'-diiodo analog were amenable to unilateral but only to a double-sided halogen/metal permutation. 874907-53-69 RE: SPN (Synthetic preparation); PREP (Preparation) (preparation of difluoro-2,2'-binaphthalene dicarboxylic acid and study of reaction of (fluoro)naphthalene and atropisomer derivative dibromo-difluoro-binaphthalene with alkyllithium compds.) 874907-53-6 CA [2,2'-Binaphthalene]-3,3'-dicarboxylic acid. 1,1'-difluoro- (9CI) (CA INDEX NAME)

THERE ARE 34 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE

L4 ANSWER 2 OF 11 CA COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 143:248127 CA Efficient optical resolution of secondary alkyl alcohols by chiral supramolecular hosts

AUTHOR(S): Inai, Yoshitane; Sato, Tomohiro; Kuroda, Reiko

SURCE: Meguro-ku, 153-0041, Japan

Chemical Communications (Cambridge, United Kingdom) (2005), (26), 3289-3291

CODEN: CHOOPS; ISSN: 1359-7345

PUBLISHER: Royal Society of Chemistry

Journal

DOCUMENT TYPE: LANGUAGE:

LOGAL 178: DOLLARS Register Re solution

solution

Due to rotation, [1,1'-biphenyl]-2,2'-dicarboxylic acid (I) is not chiral
in solution; however, in a complex with (1R,2R)-1,2-diphenyl-1,2ethanediamine (II), this compound can exhibit axial chirality. When a

ethanediamine (II), this compound can exhibit axial chirality. When a solution of I, II, and racemic butanol was mixed, a I-II host-guest compound was formed, wherein (\$)-2-butanol was trapped between a hydrogen bond between the hydroxyl group and biphenyl acid anion. The conformation of I was fixed to be axially chiral, (R)-I.

IT 2178-01-2

2178-03-2
RL: PEP (Physical, engineering or chemical process); PYP (Physical process); PROC (Process)
(optical resolution of secondary alkyl alc. derivs. via formation of supramol. multi-chiral inclusion complexes from binephthalenedicarboxylic acid-(R,R)-di(phenyl)ethanediamine-chiral

alc.)
2178-03-2 CA
(2,2'-Binsphthalene]-3,3'-dicarboxylic acid (7CI, 8CI, 9CI) (CA INDEX NAME)

REFERENCE COUNT: THIS

23 THERE ARE 23 CITED REFERENCES AVAILABLE FOR

FORMAT

RECORD. ALL CITATIONS AVAILABLE IN THE RE

(Continued)

L4 ANSWER 1 OF 11 CA COPYRIGHT 2006 ACS on STN

ANSWER 3 OF 11 CA COPYRIGHT 2006 ACS on STN ESSION NUMBER: 141:260931 CA

L4 ANSWER 3 OF 11 CA
ACCESSION NUMBER:
1111E:
111260931 CA
Advanced Method for Assignment of Absolute
Configuration Utilizing an Induced CD and
Computational Technique: Its Application to Natural
Products Possessing a Secondary Alcohol
Hosoi, Shinzo; Serata, Jun; Kluchi, Pumiyuki;
Sakushima, Akiyo; Ohta, Tomihisa
SCHOOl of Phermaceutical Sciences, Kyushu University
of Health and Welfare, Nobeoka, 882-8568, Japan
Journal of Natural Products (2004), 97(9), 1568-1570
COBEN: JNPRDF; ISSN: 0163-1664
American Chemical Society
Journal
AB A modified procedure for determining absolute configurations using an
induced CD
method and mol. mechanics calcus. is disclosed. The practical usefulness
of the present technique was demonstrated by its application to a few
natural products.

11 385707-15-9
RL: RCT (Reactant); RACT (Reactant or reagent)
(sesignment of absolute configuration of natural products possessing a
secondary alc. Utilizing an induced CD and mol. mechanics calcus.

9
binaphthalene derivative)
386707-15-9 CA
[2,2'-Binaphthalene]-3-carboxylic acid, 3'-(cyanocarbonyl)-, methyl ester
(9CI) (CA INDEX NAME) Gad Desch

REFERENCE COUNT:

THERE ARE 18 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L4 ANSWER 4 OF 11 CA ACCESSION NUMBER: TITLE:

COPYRIGHT 2006 ACS on STN
138:73974 CA
Preparation of achiral biaryl-type compounds, their
use as chromophores for circular dichroism (CD), and
determination of absolute configuration of chiral
compounds
Ota, Tomihisa, Hosoi, Shinzo
Kanazawa University, Japan
Jpn. Kokai Tokkyo Koho, 18 pp.
CODEN: JKXXAF
Patent
Japanese 1

INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
JP 2003002871	A2	20030108	JP 2001-167770		20010621
US 2003088104	A1	20030508	US 2002-82251		20020226
US 6727098	B2	20040427			
US 2004171662	A1	20040902	US 2004-785070		20040225
PRIORITY APPLN. INFO.:			JP 2001-187770	λ	20010621
			HE 2002-82251		20020226

OTHER SOURCE(S): MARPAT 138:72974

AB Determination of absolute configuration of chiral alcs., thiols, or amines involves introduction of achiral biaryl compds. I (R = H, He, Et, iso-Pr, n-Bu, iso-Bu, tert-butyl; X = H, Me, Me2N, Me0, NO2, NH2, CN, CI, Br; Y = CH, CN, indid20:1-1:yl, 1,3,4-tris20:1-yl; when R = H, Y = OH, then X = Me2N, CN; when R = Me, Y = OH, then X = Me, Me2N, NO2, NH2, CN; when R = Et, Y

OH, then X = Me, Me2N, Me0, NO2; X = H, Y = OH, then R = tert-butyl) or their analogs as CD chromophores to the chiral compds. and, is based on the relative bulk of the substituents in the α C, the priority in the CIP method, and the exciton chirality. Thus, 1- or d-menthol was esterified with 3-cyanocarbonyl-3'-methoxycarbonyl-2,2'-binaphthalene in the presence of DMAP to give (R)- or (S)-ester, resp. Their exciton chirality was - and +, resp. 106653-99-0, 3-Carboxy-3'-methoxycarbonyl-2,2'-binaphthalene

ANSWER 4 OF 11 CA COPYRIGHT 2006 ACS on STN (Continued)

482359-73-9 CA [2,2'-Binaphthalene]-3,3'-dicarboxylic acid, mono(2-methylpropyl) ester (SCI) (CA INDEX NAME)

386707-15-9P, 3-Cyanocarbonyl-3'-methoxycarbonyl-2,2'-

binaphthalene RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation of achiral biaryl-type compds. as CD chromophores for determination of

absolute configuration of chiral compds.)
386707-15-9 CA

(9CI) (CA INDEX NAME)

482359-70-6 CA [2,2'-Binaphthalene]-3,3'-dicerboxylic acid, mono(1-methylethyl) ester (9C1) (CA INDEX NAME)

482359-71-7 CA
[2,2'-Binaphthalene]-3,3'-dicarboxylic acid, monobutyl ester (9CI) (CA
RMDEX NAME) RN CN

482359-72-8 CA [2,2'-Binephthalene]-3,3'-dicarboxylic acid, mono(1,1-dimethylethyl)

(9CI) (CA INDEX NAME)

L4 ANSWER 5 OF 11 CA COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 137:185699 CA
TITLE: A Biomimetic Transformation of Serratinine into
Serratezomine A through a Modified Polonovaki

Reaction AUTHOR(S): CORPORATE SOURCE:

Morita, Hiroshi; Kobayashi, Jun'ichi Graduate School of Pharmaceutical Sciences, Hokkaido University, Sapporo, 060-0812, Japan Journal of Organic Chemistry (2002), 67(15),

SOURCE: 5378-5381

CODEN: JOCEAH; ISSN: 0022-3263 American Chemical Society Journal

PUBLISHER: DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S):

LANGUAGE: Journal
LANGUAGE: Beglish
OTHER SOURCE(S): CASREACT 137:185699

AB Application of a modified Polonoveki reaction for serratinine resulted in generation of serratezomine A with a novel seco-serratinine-type skeleton recently isolated from the club moss Lycopodium serratum var. serratum. This biomimetic transformation supports a biogenetic pathway proposed for serratezomine A. The absolute stereochem. of serratezomine A was established
by an induced context of the context o

bismed by an induced exciton chirality and modified Mosher methods. 386707-15-9

385707-15-9
RL: RCT (Reactant); RACT (Reactant or reagent)
(biomimetic transformation of serratinine into serratezomine A through a modified Polonovski reaction)
385707-15-9
CA [2,2'-Binsphthalene]-3-carboxylic acid, 3'-(cyanocarbonyl)-, methyl ester
(SCI) (CA INDEX NAME)

Ond Date

21

THERE ARE 21 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE

REFERENCE COUNT: THIS FORMAT

L4 ANSWER 6 OF 11 CA COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 136:85715 CA TITLE: Novel development of exci 136:85715 CA
Novel development of exciton-coupled circular
dichroism based on induced axial chirality
Hosoi, Shinzo; Kamiya, Makiko; Ohta, Tomihisa
Faculty of Pharmaceutical Sciences, Kanazawa
University, Kanazawa, 920-0934, Japan
Organic Letters (2001), 3(23), 3659-3662
CODEN: ORLEP7; ISSN: 1523-7060
American Chemical Society
Journal AUTHOR (S) : CORPORATE SOURCE: SOURCE: PUBLISHER: LANGUAGE: English
OTHER SOURCE(S): CASREACT 136:85715
AB A simple method for determining the absolute configuration of chiral sics. with a . with a unique chromophoric reagent, 3-cyanocarbonyl-3'-methoxycarbonyl-2,2'-binaphthalene (I), based on induced exciton chirality has been develope Thus, the alcs. were reacted with I to give the esters. The UV and CD data was collected. The structural feature of the a-positions of the carbinol carbon was found to be important to correlate the sign of developed. Cotton effect and the absolute stereochem. of the alcs. Practical usefulness of the present method was demonstrated by the determination of the absolute configuration of 17,18-dihydroxybergamottin. RL: RCT (Reactant); RACT (Reactant or reagent) (absolute configuration of chiral alcs. via UV and exciton-coupled CD binaphthalene derivative)
106653-99-0 CA
(2,2'-Binaphthalene)-3,3'-dicarboxylic acid, monomethyl ester (9CI) (CA
INDEX NAME) 66222 386707-15-99 RE: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (abbolute configuration of chiral alcs. via UV and exciton-coupled CD binaphthalene derivative)
385707-15-9 CA
(2.2'-Binaphthalene]-3-carboxylic acid, 3'-(cyanocarbonyl)-, methyl ester
(SCI) (CAINDEX NAME) L4 ANSWER 7 OP 11 CA COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 67:90570 CA COMPRIGHT 2006 ACS on STN
ACCESSION NUMBER: 67:90570 CA COMPRIGHT 2006 ACS on STN
ACCESSION NUMBER: 67:90570 CA COMPRIGHT 2006 ACS on STN
ACCESSION NUMBER: 7:90570 CA
AUTHOR(S): 05 pollycyclic aromatics. XIV. Ozonation of pentaphene and benzo[rat]pentaphene
AUTHOR(S): Fordham Univ., New York, NY, USA
JOURNAI OF OFMAM UNIV., NEW YORK, NY, USA
JOURNAI OFMAM UNI XIV. Ozonation oxidation to
cis-6,7-dihydroxy-6,7-dihydropentaphene followed by aqueous NaIO4
oxidation
oxidation
oxidation of II gave 14% IV, while IV was independent Chromic acid oxidation of II gave 14% IV, while IV was independently Chromic acid Oxidation of II gave 14% IV, while IV was independently ared in 718 yield via Cu20 coupling of the diszonium salt of 2-aminonaphthalene-3-carboxylic acid. II in base underwent an intramol. Cannizzaro reaction to 2,2°-binaphthyl-3-hydroxymethyl-3'-carboxylic acid which lactonized on treatment with strong acid or mild heat to an e-lactone. Ozonolysis of I with 4 mole equive. O3 followed by oxidative work-up gave 9% III and 53% 2,2',4,4',5,5'-hexacarboxybiphenyl (V). The hexa-Me ester obtained from V was independently synthesized by an Ullman coupling of 5-bromo-1,2,4-tricarbomethoxybenzene. Ozonization of benzo(rst)pentaphene (VI) in CH2Cl2 at -78° with 3.5 mole equivalent O3, followed by oxidative work-up led to 17% benzo(rst)pentaphene-5,8-dione, 4% III, 10% p-terphenyl-3,2',3'-tetreacrboxylic acid 2',3'-anhydride, and 3% 2-(o-carboxyphenyl)-1,10-phenanthrenedicarboxylic acid anhydride, with a 56% recovery of VI. A comparison of the tivity prepared reactivity
to G3 of the noncarcinogenic I and related pentacyclic and hexacyclic
hydrocarbons of increasing carcinogenicity indicates that there is no
simple, consistent correlation between carcinogenicity, K- and L-region
additivity toward G3, and the Pullman (P. and P. CA 50. 4756b) electronic theory of carcinogenesis. 35 references. 2178-03-2P RL. SPN (Synthetic preparation); PREP (Preparation) (preparation of) 2178-03-2 CA 2178-03-2 CA [2,2'-Binaphthalene]-3,3'-dicarboxylic acid (7CI, 8CI, 9CI) (CA INDEX NAME) G100,20

ANSWER 6 OF 11 CA COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT: THIS

FORMAT

21 THERE ARE 21 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE

L4 ANSWER 8 OF 11 CA COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
ORIGINAL REFERENCE NO.:
CITIES:
AUTHOR(S):
CORPORATE SOURCE:
SOURCE:
DOCUMENT TYPE:
DOCUMEN

reaction
on 11H-benzo[b]fluoren-11-one oxime (22% total). Reduction of the

on In-Denzolbiluoren-il-one oxime [224 Cotal]. Reduction of the same with lithium alumium hydride gave the 5.6-dihydrobenzo[b] and -[j]phenanthridines. Dehydrogenation of these dihydro derivatives produced the parent aromatic heterocycles benzo[b] - and benzo[l]phenanthridine in best overall yields of 20% and 12%, respectively. A few substituted benzophenanthridines were also prepared. Assignment of structures was based on uv, ir, and N.M.R. spectrs of the dihydro derivatives as well as on separate unequivocal synthesis of the isomeric benzophenanthridines.
2178-03-2, [2,2'-Binaphthalene]-3,3'-dicarboxylic scid (preparation of) 2178-03-2 CA [2,2'-Binaphthalene]-3,3'-dicarboxylic acid (7CI, SCI, 9CI) (CA INDEX NAME)

IT





Page 5

OTHER SOURCE(S)

JUNIONI TYPE: Journal
JUNGE: Unavailable
RE SOURCE(5): CASREACT 58:53289
For diagram(s), see printed CA Issue.
cf. CA 52, 20099f. Reactions which normally result in nuclear coupling
led to reductive dehalogenation of 2-substituted 1-halomaphthalenes,
except in the case of Me 1-bromo-2-naphthoate, which, by an Ullmann
reaction and further steps, was converted into 1,1'-binaphthyl-2,2'dialdehyde. Starting with the nuclear coupling of diazotized
3-amino-2-naphthoic acid, a similar synthesis of 2,2'-binaphthyl-3,3'dialdehyde was carried out. Reaction of hydrazine with the former
dialdehyde was carried out. Reaction of hydrazine with the former
dialdehyde underwent reductive cyclization exclusively, giving pentaphene
(II) in 40% overall yield from the amino acid.
90115-51-6, [2,2'-Binaphthalene]-3,3'-dicarboxylic acid, ethyl
ester 106653-99-0, [2,2'-Binaphthalene]-3,3'-dicarboxylic acid, ethyl
ester (preparation of)
90135-51-6 CA
(2,2'-Binaphthalene]-3,3'-dicarboxylic acid, ethyl ester (7CI) (CA INDEX
NAME)

106653-99-0 CA [2,2'-Binaphthalene]-3,3'-dicarboxylic acid, monomethyl ester (9CI) (CA KNDEX NAME)

L4 ANSWER 10 OF 11 CA COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:
36:2769 CA

ORIGINAL REFERENCE NO.: 36:4461,447a-g
Polycyclic aromatic hydrocarbons. XXVIII.
Dibenzofluorenes

AUTHOR(S):
Martin, Richard H.
Journal of the Chemical Society (1941) 679-85

CODEN: JCSOA9; ISSN: 0368-1769

DOCUMENT TYPE:
Journal
LANGUAGE:
Unavailable
GI For diagram(s), see printed CA Issue.
AB cf. C. A. 35, 4008.3. Heating 25 g. of 3,2-BrC10H6CO2Me (bl.5
170°) with 18 g. Cu bronze at 190-200° gives 18.8 g. of the
Me ester, b0.6 260-70°, m. 173-3.5°, of 2,2'-binaphthyl-3,3'dicarboxylic acid (I), m. 298-9°. I could not be converted into
2,3,6,7-diabenzofluorenone (II) by boiling with Ac20; heating 5 g. of the
Pb salt a 0.2 mm. over a free flame gives 1.5 g. NNM4 MO at

269-70°. Reduction of 1 g. of II by heating with 6 cc. N2H4.H2O at 255° for 8 hrs. gives 2,3,6,7-dibenzofluorene, m. 282.5-3.5°, sublimes at 210°/0.1 mm. Heating 0.5 g. of II with 3 g. KOH for 0.5 hr. at 240-50° gives 2,2'-binephthyl-3-carboxylic acid, m. 189-91°; heating with 50 parts of 80° H3SO4 on the water bath for 3 hrs. gives 1,2,6,7-dibenzofluorenne (IIA), orange, m. 211°; concentrated H2SO4 gives a carmine-red solution 1,2-BrC1OH6CO2Me (19.7 g.) and 3 g. Cu bronze, heated at 190° and 10 g. of the Cu added in portions during 0.5 hr., with heating for an addnl. 4.5 hr.,

added in portions during 0.5 hr., with heating for an addnl. 4.5 hr.,

8.5 g. of Me 1,1'-binephthyl-2,2'-dicarboxylate, m. 156.5-7.5'; the
free acid (17.4 g.), refluxed 0.5 hr. with excess of Ac20 and the residue
heated at 280° for 3 hrs., gives 9.4 g. of 3,4.5.6dibenzofluorenone (III), dark red, m. 222-2.5'; the H2504 solution is
carmine-red, oxime, oxange-red, m. 253-4'. Reduction of 2 g. of
III with N3H4.H20 (15 hrs. at 180') gives 1.35 g. of
3,4.5,6-dibenzofluorene (IV), m. 152-2.5'; dipicrate, reddish
brown, m. 154-5', oxidation of IV with Se02 gives III. Pusion of
III with Alkil3-Nacl gives 1.2,8,9-dibenzathrone, yellow, m.
185-6'. 1,2,7,8-Dibenzofluorene (V) (1.6 g.) with Se02 at
230° for 6 hrs. gives 1.2 g. of 1,2,7,8-dibenzofluorenom (VI), m.
263-5.5'; fusion with K0H at 240-50' gives
2,2'-binsphthyl-1-carboxylic acid, m. 177-9', which with 804 H2504
at 100° for 4 hrs. gives VI; reduction of VI with N3H4.H20 yields
V. This behavior, together with the synthesis of II and IIA, establishes
the structure of V and VI. 1-C10H7COC1 (50 g.) and 36 g. tetralin in 40
cc. C52, added to 38 g. of Alc13 in 80 cc. C52 in an ice bath, give 44 g.
of the Ketone, bo.8 230-5' (oxime, C2H1H8ON, m. 172-2.5');
dehydrogenation with 5 at 220' yields 20 g. of 1,2'-dinaphthyl
Atone, 5 g. of which is reduced by Amonk to 4 g. of the carbinol (VII).
Cyclization of 3 g. of VII by 6 g. HPO3 gives 1,2,5,6-dibenzofluorene but
the yield is too small for the method to be of practical use. Reaction
chloromethyltetralin (32.9 g., b20 148') with 31.8 g. McCH(CO2Et)2

chloromethyltetralin (32.9 g., b20 148°) with 31.8 g. MeCH(COZEt)2 and 4.2 g. Na in 120 cc. C6H6 gives 29.5 g. of the ester. C19H2604, b0.4 160-1°, heating the acid at 170° gives 23.8 g. of 6-tetrallyl-e-methylpropionic acid, b0.1 15°; the acid chloride with SnCl4 in C6H6 gives a mixture of ketones (VIII and IX),

125-35°; about 10% crystallized from petr. ether at -2° and m. 80.5-1.5°; the liquid b0.1 123°. Oxidation of the ketones gives only mellophanic acid. Reaction of the ketones with PhCH2CH2MgCl

L4 ANSWER 9 OF 11 CA COPYRIGHT 2006 ACS on STN (Continued)

ANSWER 10 OF 11 CA COPYRIGHT 2006 ACS on STN (Continued) and dehydration of the carbinol with KHS04 gives a hydrocarbon, C22H24, b0.1 174*; cyclization with AlCl3 in CS2 gives the satd. isomer, b0.15 176*; Se at 305* gives a hydrocarbon, C21H14, m. 306-8*.

2178-03-2, [2,2'-Binaphthalene]-3,3'-dicarboxylic acid (and derivs.)

2178-03-2 CA [2,2'-Binaphthalene]-3,3'-dicarboxylic acid (7CI, 8CI, 9CI) (CA INDEX NAME)

LA ANSWER 11 OF 11 CA COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 22:22028 CA ORIGINAL REFERENCE NO.: 22:2572d-e Naphthalene derivatives HAPPER ASSIGNEE(S): DOCUMENT TYPE: LANGUAGE: PATENT INFORMATION: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

OB 278100 19271006 GB 1926-16931 19260706

AB Dinaphthyldicarboxylic acids and their substitution products are obtained by treating diato compds. derived from o- or peri-mainonaphthoic acids or their derivs. with suitable reducing agents such as an ammoniacal solution of Cu20 or a neutral solution of Na2603 or a ferrous salt. Examples are given of the production of 1,1'-dinaphthyl-8,8'-di-carboxylic acid, 2,2'-dinaphthyl-3,2'-dicarboxylic acid didethyl seter.

1,1'-dichloro-2,2'-dinaphthyl-3,3'-dicarboxylic acid dethyl seter.

1,1'-dichloro-2,2'-dinaphthyl-3,2'-dicarboxylic acid, 4,4'-didhoro-1,1'-dinaphthyl-8,8'-dicarboxylic acid, 4,4'-didhoro-1,1'-dinaphthyl-8,8'-dicarboxylic acid, 5,5'-dimethoxy-1,1'-dinaphthyl-8,8'-dicarboxylic acid and the corresponding diethoxy compound Cf. C. A. 23, 2380.

IT 2178-03-2, (2,2'-Binaphthalene]-3,3'-dicarboxylic acid, 1,1'-dichloro-(preparation of)
RN 2178-03-2 CA

RN 2178-03-2 CA

RAME)

RN 859931-11-6 CA CN [2,2'-Binaphthalene]-3,3'-dicerboxylic acid, 1,1'-dichloro- (3CI) (CA INDEX RNME)

L4 ANSWER 11 OF 11 CA COPYRIGHT 2006 ACS on STN (Continued)

B407129

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10/785,070
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=> file marpat

=> s l1 full L5 10 SEA SSS FUL L1

=> s 15/com L6 9 L5/COM

=> d ibib abs fqhit 1-9

```
L6 ANSWER 1 OF 9 MARPAT COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 143:360057 MARPAT
ITILE: 11-beta hydroxysteroid dehydrogenase type 1
inhibitors
agents for the treatment of cancers, especiall

cancer Vander Jagt, David L.; Royer, Robert E.; Deck, Lorraine M.

PATENT ASSIGNEE(S): USA
SOURCE: U.S. Pat. Appl. Publ., 15 pp.
CODEN: USAXCO
DOCUMENT TYPE: Patent
LANGUAGE: PARILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.
                                                                           as anti-obesity/anti-diabetes compounds and 17-beta
hydrosteroid dehydrogenase type i inhibitors as
                                                                           agents for the treatment of cancers, especially
                                                                                                                                                                                                                                                                   Patent location:
Note:
                                                                  KIND DATE
A1 20051013
                                                                                                                  APPLICATION NO. DATE

US 2005-93493 20050330

US 2004-560387P 20040408

Biscovery that 11-Beta Hydroxysteroid don mol. etiol. for visceral obesity and as well a treatment for diabetes,
    PRIORITY APPLN. INFO.:

AB This invention is directed to the discovery that 11-Beta Hydroxystero:

Dehydrogenase Type 1 may be a compon mol. stiol. for viaceral chesity the metabolic syndrome of obesity as well a treatment for diabetes, especially type II diabetes. The present invention also relates to the use of certain compds. as inhibitors of 17-Beta Hydroxysteroid Dehydrogenase Type
      Type

1 and their use for the treatment of cancer, especially breast cancer.
      16 (0)-G2
    L6 ANSWER 2 OF 9
ACCESSION NUMBER:
IIILE:

MARPAT COPYRIGHT 2006 ACS on STN
138:72974 MARPAT
Preparation of achiral biaryl-type compounds, their
use as chromophores for circular dichroism (CD), and
determination of absolute configuration of chiral
                                                                           occementation of absolute cont
compounds
Ota, Tomihisa; Hosoi, Shinzo
Kanazawa University, Japan
Jpn. Kokai Tokkyo Koho, 18 pp.
CODEN: JKXXAF
     INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:
      DOCUMENT TYPE:
                                                                           Patent
Japanese
     LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                                                                                                                                                                                                                                                  G1 = OH
G4 = OH
Patent location:
                    PATENT NO.
                                                                  KIND / DATE
                                                                                                                              APPLICATION NO. DATE
                                                                                 20030108
20030508
20040427
20040902
                   JP 2003002871
                                                                    A2
A1
B2
                                                                                                                             JP 2001-187770
US 2002-82251
                                                                                                                                                                             20010621
     US 2003088104
US 6727098
US 2004171662
PRIORITY APPLN. INFO.:
                                                                                                                             US 2004-785070
JP 2001-187770
US 2002-82251
                                                                                                                                                                              20040225
     GΙ
    AB Determination of absolute configuration of contents of contents of amines involves introduction of achiral biaryl compds. I (R = H, Me, Et, iso-Pr, n-Bu, iso-Bu, tert-butyl; X = H, Me, Mean, Meo, NO2, NH2, CN, Cl, Br; Y = CH, CN, imidazol-1-yl, 1,3,4-triazol-1-yl; when R = H, Y = OH, then X = Me2N, NO2, NH2, CN; when R = Et, Y CN; when R = Me, Y = OH, then X = Me, Me2N, NO2, NH2, CN; when R = Et, Y
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OH, then X = Me, Me2N, MeO, NO2; X = H, Y = OH, then R = tert-butyl) or their analogs as CD chromophores to the chiral compds. and, is based on the relative bulk of the substituents in the α C, the priority in the CIP method, and the exciton chirality. Thus, 1 or d-menthol was esterified with 3-cyanocarbonyl-3'-methoxycarbonyl-2,2'-binaphthalene in the presence of DMAP to give $\{R\}$ - or $\{S\}$ -ester, resp. Their exciton chirality was - and +, resp.

L6 ANSWER 1 OF 9 MARPAT COPYRIGHT 2006 ACS on STN (Continued) claim 1 and pharmaceutically acceptable salts ANSWER 2 OF 9 MARPAT COPYRIGHT 2006 ACS on STN (Continued) claim 1 substitution is restricted

METR 2

L6 ANSWER 3 OF 9
ACCESSION NUMBER:
136:387542 MARPAT
TITLE:
Black waterborne storage-stable ink-jet inks and
printing method using them
Adachi, Keiichi
Puji Photo Pilm Co., Ltd., Japan
JDN. Kokai Tokkyo Koho, 19 pp.
CODENT TYPE:
DOCUMENT TYPE:
Patent
AMININGE.
JENERAL JENERAL
ANGELINGE.
JENERAL JENERAL
ANGELINGE.
JENERAL JENERAL
ANGELINGE.
JENERAL JENERAL
ANGELINGE. DOCUMENT TYPE: LANGUAGE: PAMILY ACC. NUM. COUNT: PATENT INFORMATION: KIND DATE PATENT NO. APPLICATION NO. DATE JP 2002146249 A2 2082052 JP 2000-341683 JP 2000-341683 20001109 PRIORITY APPLN. INFO.: I The inks contain a dye of I type (R1-3 = H, halogens, alkyl, aryl, CH, acyl, carbamoyl, alkoxycarbonyl, aryloxycarbonyl, acyloxy, alkoxy, alkoxy, aryloxy, alkylution, arylatio, arylatio, aulfamoyl, alkylutionyl, arylaulfonyl or amino groups; R4, R5 = H, alkyl, aryl groups; R6-9 = H, halogens, alkyl, aryl, carbamoyl, alkoxy, arylathio, arylfatho, sulfamoyl, alkylaulfonyl, arylsulfonyl or amino groups provided that at least 1 of to R9 is sulfonic acid or carboxylic acid or their salts), optionally a naphthalene type azo dye and other additives. MSTR 2 G1---G6 G1 - naphthyl (substd. by 1 or more G2)
G2 - CO2H
G6 - naphthyl (substd. by 1 or more G2)
Patent location: claim 3 L6 ANSWER 4 OF 9
ACCESSION NUMBER: 13:223160 MARPAT
TITLE: Epoxides with a liquid crystalline phase, and processes and photoinitiators for their conversion to epoxy resins
INVENTOR(S): Schnutfeil, Guenter; Schroeder, Hendrik; Hartwich, Andreas Experiment assignment of the conversion to epoxy resins
Schnutfeil, Guenter; Schroeder, Hendrik; Hartwich, Andreas Experiment assignment of the conversion to epoxy resins
FATENT ASSIGNME(S): Praunhofer-Gesellachaft zur Foerderung der PATENT ASSIGNEE(S): Angewandten Porschung e.V., Germany Ger. Offen., 13 pp. CODEN: GWXXBX Patent German SOURCE: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: KIND DATE PATENT NO. APPLICATION NO. DATE DE 10004442 PRIORITY APPLN. INFO.: DE 2000-10004442 20000202 DE 1999-19904028 19990202 A1 20000907 - (CH₂) n1Y1Z1XZY (CH₂) n The epoxides have the general formula I $\{R = H, ary\}$, alkyl; R1 = H, alkyl; X = direct link, CO2, CR2:CR2, CR2:N, $CH:N\{O\}$, N:N, $N:N\{O\}$; each - H, alkyl; Y, Y1 = 0, S, CH2, CO2; Z, Z1 = (un)substituted divalent aromatic, alicyclic, or heterocyclic group; n, n1 = 1-16). The photoinitiator for manufacture of the epoxy resin is selected from imidazole derivs., BF3 complexes, Fe(II) aromatic complexes, iodonium salts, ammonium

salts and sulfonium salts. Thus, 4-HOC6H4CO2C6H4OH-4 was etherified with
2 equiv Br(CH2)4CH:CH2, and the product was epoxidized with 3-ClC6H4CO2OH
to give a diepoxide which exhibited a liquid crystalline phase between and 54°. MSTR 18

ANSWER 4 OF 9 MARPAT COPYRIGHT 2006 ACS on STN (Continued) Patent location: claim 1

ANSWER 3 OF 9 MARPAT COPYRIGHT 2006 ACS on STN substitution is restricted

(Continued)

- 82-5 87-8

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L6 ANSWER 5 OF 9 MARPAT COPYRIGHT 2006 ACS ON STN

ACCESSION NUMBER:
1111E: 126:199271 MARPAT
Process for the hydroformylation of water-insoluble unsaturated compounds with rhodium-phosphine catalyst systems
Bahrmann, Helmut; Lappe, Peter; Fell, Bernhard; Xia, Zhigao; Kanagasabapathy, Subba
HOGCHAE A.-G., Germany
Ger. Offen., 10 pp.
CODENT TYPE: Patent MARPAT
PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

DE 19521393 Al 19970306 DE 1995-19532393 19950902
TW 349751 B 20000511 TW 1996-85109975 19960816
CA 2184048 AA 19970301 CA 1996-2184048 19960823
CA 2184048 C 19990525
EP 761635 Bl 19990804
R: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE

AT 182876 E 19990815 AT 1996-113605 19960824
CA 9643704 T3 19991216 ES 1996-113605 19960824
CA 9664390 A1 19970301 CA 1996-2728 19960826
DL 1986-21364 T3 19991316 ES 1996-113605 19960824
AU 9664390 A1 19970306 AU 1996-64390 19960826
AU 9664390 A1 19970301 AU 1996-64390 19960830
CN 1149041 AA 19970507 CN 1996-613503 19960830
CN 1149041 AA 19970507 CN 1996-6131503 19960830
CN 1149043 A 19970507 CN 1996-61390 19960830
CN 1149043 A 19970507 CN 1996-61390 19960830
CN 1149041 AA 19970507 CN 1996-61390 19960830
CN 1149043 A 19970507 CN 1996-61390 19960830
CN 1149043 A 19970507 CN 1996-61390 19960830
CN 1149041 AA 19970507 CN 1996-61390 19960830
CN 1149043 A 19970507 CN 1996-61390 19960830
CN 1149043 A 19970507 CN 1996-61390 19960830
CN 1149041 AA 19970507 CN 1996-61390 19960830
CN 1149041 AA 19970507 CN 1996-61390 19960830
CN 1149043 A 19970507 CN 1996-113603 19960830
CN 1149041 AA 19970507 CN 1996-113603 19960830
CN 1149043 A 19970507 CN 1996-113603 19960830
CN 1149041 AA 19970507 CN 1996-113603 19960830
CN 1149041 AA 19970507 CN 1996-11973 19960830
CN 114
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L6 ANSWER 6 OF 9 MARPAT COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 121:107982 MARPAT
ITILE: Processe for the preparation of aldehydes by
hydroformylation using rhodium-phosphine catalysts
                                                      phosphonium salt solubilizers
                                                     phosphonium salt solubilizers
Bahrmann, Helmut; Lappe, Peter
Hoechst A.-G., Germany
Eur. Pat. Appl., 10 pp.
CODEN: EPXXDW
Patent
German
  INVENTOR(S):
 PATENT ASSIGNEE(S):
SOURCE:
 DOCUMENT TYPE:
  LANGUAGE:
 LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
           PATENT NO.
                                                KIND DATE
                                                                                            APPLICATION NO. DATE
A1 19940622
B1 19961113
           CO and H2 in an aqueous solution of a water-soluble Rh-phosphine-complex
CO and H2 in an aqueous solution of a water-soluble Rh-phosphine-complex catalyst and a quaternary phosphonium salt serving as a solubilizing agent. For example, a catalyst solution was prepared from tri-Na tris(m-sulfophenyl)phosphine, tetradecyltriethylphosphonium bromide (I), H2O, buffer solution (pH 6.0), and Rh acetate, the mixture of which was heated under
           ad under synthesis gas (CO/H2 = 1:1) at 110^{\circ} and 2.5 MPa. Hydroformylation of 1-tetradecene in the catalyst solution under the same conditions for
           gave 74.2% conversion to aldehydes, with activity (mol aldehyde/mol Rh·min) of 3.10 and productivity (g aldehyde/mi catalyst solution-h) of 0.075. In contrast, a run without I gave only 0.10% conversion, with both activity and productivity values of 0.00.
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METE 3

- 22

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MARPAT COPYRIGHT 2006 ACS on STN

121:107978 MARPAT
Proparation of higher primary alcohols
Bahrwann, Helmut; Deckers, Gregor; Greb, Molfgang;
Heymannas, Peter; Lappe, Peter; Hueller, Thomas;
Szameitat, Juergen; Mielbus, Brnat
Hoechet A.-G., Germany
Eur. Pat. Appl., 7 pp.
CODEN: BEXXLDM
Patent
German

DUNT: 1
 L6 ANSWER 7 OF 9
ACCESSION NUMBER:
  TITLE:
INVENTOR(S):
  PATENT ASSIGNEE(S):
SOURCE:
  DOCUMENT TYPE:
LANGUAGE:
PAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
               PATENT NO.
                                                              KIND DATE
                                                                                                                     APPLICATION NO. DATE
PATENT NO. KIND DATE APPLICATION NO. DATA

EP 602442 A2 19940622 EP 1993-119242 19931130

R: BE, DE, ES, FR, GB, IT, NL, SE

DE 4247275 A1 19940623 DE 1992-4242725 19921217

US 6051743 A 20000418 US 1993-163086 19931207

CA 2111026 AA 19940618 CA 1993-211026 19931207

JP 06279334 A2 19941004 JP 1993-309456 19931209

JP 07039362 B4 19950501

BR 9305007 A 19940705 BR 1993-5007 19931210

AU 9305292 A 19940804 CA 1993-292 19931210

AU 93052439 A1 19940804 CA 1993-52439 19931215

AU 664126 B2 19951102

PRIORITY APPLN. INFO:

DE 1992-4242725 19921217

AB The title process comprises hydroformylation of a (Pischer-Tropsch)

olefin
 AB T
              in the presence of Rh or a compound thereof, a water-soluble phosphine,
              salt comprising a Z+ABCD cation [A = (ar)alkyl; B,C,D = alkyl; Z = N or
              and a water-soluble sulfonated or carboxylated aromatic phosphine anion
              by hydrogenation. Thus, a mixture comprising a primarily
 nonene-containing
Pischer-Tropsch olefin, a water solution of [3-(NaO3s)C6H4]3P and the
corresponding trimethyltetradecylammonium salt, Rh acetate, and a
NaOac/HOAc buffer was maintained 6h at 125° under 2.5MPa CO/H to
give 85 olefin conversion.
       MSTR 1
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L6 ANSMER 8 OF 9
ACCESSION NUMBER: 120:134812 MARPAT
TITLE: Preparation of methylenediphosphonic acid derivatives as drugs
INVENTOR(S): Tanahashi, Masahiko; Senba, Yuriko; Nakadate, Akio; Kawabe, Norio; Uchiro, Takumi
TOrey Industriee, Japan
Jpn. Kokai Tokkyo Koho, 15 pp.
COLINIT TYPE: Patent DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION: A2 19930803 B2 20021105 AA 19940120 A1 19940120 PATENT NO. APPLICATION NO. DATE JP 05194565 JP 3341303 CA 2111670 JP 1992-183866 19920710 OTHER SOURCE(S):

(CH₂) ss (CH₂) .

AB Title compds. I [R1-R4 = H, alkyl, cation; X, Y = substituent on the naphthyl radical such as halo, nitro, alkyl, (un)substituted amino; A = (this) (oxa) (aza) polymethylene; B = H, alkyl, amino, etc.; m = 0-3 integer;
n = 0-4 integer], useful as antiinflammatories, antirheumatics, interleukin-1 inhibitors, antioxidants, inhibitors of bone resorption, as well as drugs for bone metabolism, are prepared by reacting CH3[P(0) (OR5) (OR6)] 2

Page 12

ANSWER 7 OF 9 MARPAT COPYRIGHT 2006 ACS on STN (Continued) = (0-5) CH2 = 43-19 44-36 - 89-20 90-35 claim 1

ANSWER 8 OF 9 MARPAT COPYRIGHT 2006 ACS on STN (Continued) (R5, R6 - C1-7 alkyl) with naphthylene derivs. II (Hal - halo) or III (a 0-10 integer). A mixt. of tetra-Et methylenediphosphonate, 2,2°-dinaphthyl disulfide, and BuLi in hexane-THF was stirred at room temp. for 16 to give, after pouring into ice water and treatment with HC1,

[(2-naphthylthio)methylene]diphosphonic acid tetra-Et ester, which was
treated with Me35iCl in CH2Cl2 at room temp. for 72 h and the product
refluxed in aq. MeOH for 30 min to give

[(2-naphthylthio)methylene]diphosp
honic acid. In an in vitro study this showed 41.7% inhibition against
interleukin-1.

MSTR 3

G1 = naphthyl (opt. substd. by 1 or more G2)
G2 = CO2H
G7 = bond
Patent location: claim 3

ACCESSION NUMBER:

116:235246 MARPAT

TITLE:

Process for preparing diketones and keto acida

Malker, Theodore Roosevelt, Jr.; Jackson, Minston

Jerome, Jr.; Pleischer, Jean Carroll

PATENT ASSIGNEE(S):

Eastman Kodak Co., USA

SOURCE:

POT Int. Appl.. 30 pp.

COODEN, PIXXD2

PATENT ANDIAGE:

PATENT NO. KIND DATE

APPLICATION NO. DATE

APPLICATION NO. DATE

MI 19920206 NO 1991-US5122 19910722

MI CA, JP

RN: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LU, NL, SE

US 5107039 A 19920421 US 1990-556678 19900723

EP 540653 A1 19930512 EP 1991-141436 19910722

R; AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LU, NL, SE

JP 05508416 T2 19931125 JP 1991-513266 19910722

R; AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LU, NL, SE

PRIORITY APPLN. INFO:

US 1990-556678 19900723

WO 1991-US5122 19910722

OTHER SOURCE(S):

CASREACT 116:235246

AB The title compds. were prepared by reaction of a dicarboxylic acid and an aromatic compound in the presence of an alkanesulfonic acid and an organic and carroll beneated in the presence of MeSO1H and (CF1CO)20 to give a 95% yield of 1,3-bis(4-phenoxybenzoyl)benzene, which contained 7% tetraketone oligomer.

METE 2B

G1 = 283-1 282-3

G9 = 331-2 330-150

L6 ANSWER 9 OF 9 MARPAT COPYRIGHT 2006 ACS on STN (Continued)

Patent location: claim 1

Page 13

=> d his

(FILE 'HOME' ENTERED AT 13:37:59 ON 05 JUN 2006)

FILE 'REGISTRY' ENTERED AT 13:38:10 ON 05 JUN 2006

L1 . STRUCTURE UPLOADED

L2 0 S L1 SAM

L3 10 S L1 FULL

FILE 'CA' ENTERED AT 13:38:54 ON 05 JUN 2006

L4 11 S L3

FILE 'MARPAT' ENTERED AT 13:39:08 ON 05 JUN 2006

L5 10 S L1 FULL

L6 9 S L5/COM

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---Logging off of STN---

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Executing the logoff script...

=> LOG Y

STN INTERNATIONAL LOGOFF AT 13:39:44 ON 05 JUN 2006